Figure 1 125P5C8 SSH sequence

 Figure 2 Double stranded sequence and ORF for 125P5C8-Pro-pCR2.1.

| | M | Т | s | L | W | R | E | I | L | L | E | S | L | L | G | С | V | s |
|-------|----------|----------|-----------|----------|----------|-----------|----------|-----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|
| 1 | | | | CTG | | | GAA | | CTC P | TTG M | GAG I | TCG Y | CTG Y | CTG F | GGA P | TGT L | GTT O | TCT T |
| 55 | W TGG | S TCT | L CTC | Y TAC | H CAT | D GAC | L CTG | G GGA | | | | | TAC | | CCT | | CAA | |
| | L | E | L | T | G | L | E | G | F | S | I | A | F | L | S | P | I | F |
| 109 | CTA L | GAA T | CTC | ACT T | GGG P | F | GAA W | GGT K | TTT L | AGT V | ATA N | GCA K | K | CTT W | TCT M | L | ATA T | TTC L |
| 163 | CTA | ACA | | ACT | CCT | TTC | | | | | | AAG | | | ATG | | | |
| 217 | L | R | I ATIA | I ATC | T | I Tura | G | S | I ATA | A | S | F TTC | Q CAG | A GCT | P CCA | N TAA | A GCC | K AAA |
| 217 | L | R | L | M | V | L | A | L | G | V | s | s | S | L | I | V | Q | A |
| 271 | CTT | | | ATG | - | CTT | | CTT | | | TCT | TCC | | | ATA | | CAA | |
| 325 | V GTG | T ACT | W TGG | W TGG | S TCG | G GGA | S AGT | H CAT | L TTG | Q CAA | R AGG | Y TAC | L CTC | R AGA | I ATT | W TGG | G GGA | F TTC |
| | I | L | G | Q | I | V | L | V | V | L | R | I | W | Y | Т | s | L | N |
| 379 | ATT P | TTA | GGA W | CAG S | ATT Y | GTT Q | CTT M | GTT S | GTT N | CTA K | CGC V | ATA I | TGG L | TAT T | ACT | TCA S | CTA A | AAC I |
| 433 | | | | AGT | | - | ATG | | | | | ATA | _ | _ | | | GCC | |
| 407 | A | T | L | D | R | I | G | T | D | G | D | C | S | K | P | E | E GAA | K |
| 487 | GCC K | ACA T | G | GAT E | V | ATT | GGC T | ACA G | M | A A | S | R | AG1 P | AAA N | W | GAA L | L | AAG |
| 541 | AAG | ACT | GGT | | GTA | | | | | | | AGA | | | | | CTG | |
| 595 | G | A GCT | A GCT | F | G GGT | S AGC | L CTT | V GTG | F TTC | L CTC | T ACC | H | W TGG | V GTT | F TTT | G GGA | E GAA | V GTC |
| 373 | S | L | V | s | R | W | A | V | s | G | Н | P | Н | P | G | P | D | P |
| 649 | TCT | CTT | GTT | | AGA | | | | | | | | CAT | CCA G | GGG L | CCA M | GAT L | CCT P |
| 703 | N AAC | P CCA | F TTT | G GGA | G GGT | A GCA | V GTA | L CTG | L CTG | C TGC | L TTG | A GCA | _ | - | | ATG | _ | CCA |
| | S | С | L | W | F | R | G | T | G | L | I | W | W | V | Т | G | Т | A |
| 757 | TCT S | TGT A | TTG A | TGG G | TTT L | CGT L | GGT Y | ACT L | GGT H | TTG T | ATC W | TGG A | TGG A | GTT | ACA V | GGA S | ACA G | GCT C |
| 811 | | GCT | GCG | _ | CTC | CTT | TAC | | | | | GCA | | GCT | GTG | _ | GGC | TGT |
| 065 | V GTC | F TTC | A GCC | I | F TTT | T | A | S TCC | M | W | P | Q CAA | T | L | G GGA | H | L CTT | I ATT |
| 865 | N | S | G | T | N | P | GCA | K | T | M | т | I | A | М | I | F | Y | L |
| 919 | AAC | | | ACA | | | GGG | | | | | | | | | | TAT | CTT |
| 973 | L CTA | E GAA | I ATA | TTT T | F TTC | C TGT | A GCC | W TGG | C TGC | T ACA | A GCT | F TTT | K AAG | F TTT | V GTC | P CCA | G GGA | G GGT |
| 3.3 | V | Y | A | R | E | R | S | D | V | L | L | G | T | M | M | L | I | I |
| 1027 | GTC G | TAC L | GCT N | AGA M | GAA L | AGA F | TCA G | GAT P | GTG K | CTT | TTG N | GGG L | ACA D | ATG L | ATG L | TTA L | ATT O | ATC T |
| 1081 | GGG | | AAT | | CTA | _ | | _ | | | | CTT | | TTG | CTT | | CAA | |
| 4.105 | K | N | S | S TCT | K | V | L | F | R | K | S | E | K | Y | M | K | L | F TTT |
| 1135 | AAA L | W | AG1 L | L | V | G | V | G | AGA L | AAG L | G | L | G | L | R | Н | K | A |
| 1189 | | | | CTT | | | | | | | | | | | | | | |
| 1243 | Y TAT | E GAG | R AGA | K AAA | L CTG | G GGC | K AAA | V GTG | A GCA | P CCA | T ACC | K AAA | E GAG | V GTC | S TCT | A GCT | A GCC | I ATC |
| 1213 | W | P | F | R | F | G | Y | D | N | E | G | W | S | S | L | E | R | S |
| 1297 | TGG A | CCT H | TTC L | AGG L | TTT N | GGA E | TAT T | GAC G | AAT A | GAA D | GGG F | TGG I | TCT T | AGT I | CTA L | GAA E | AGA S | TCA D |
| 1351 | | | | CTC | | | | | | | | | | | | | | |
| | A | S | K | P | Y | M | G | N | N | D | L | T | M | W | L | G | E | K |
| 1405 | GCT L | TCT | AAG F | CCC | TAT | ATG D | GGG F | AAC G | AAT P | GAC S | TTA | ACC R | ATG Y | TGG H | T | W | GAA | AAG I |
| 1459 | TTG | GGT | TTC | TAT | ACA | GAC | TTT | GGT | CCA | AGC | ACA | AGG | TAT | CAC | ACT | TGG | GGG | ATT |
| 1513 | M ATG | A GCT | L TTG | S TCA | R AGA | Y TAC | P CCA | I ∆דיד | V GTG | K AAA | S TCT | E GAG | H | H | L CTT | L CTT | P CCG | S TCA |
| 1010 | P | E | G | E | I | Α | P | Α | I | T | L | T | V | N | I | S | G | K |
| 1567 | | | | GAG | | | | | | | | | | | | | | |
| | L | V | D | F | V | V | Т | H | F | G , | N | H | E | D | D | L | D | R |

| 1621 | CTG | GTG | GAT | TTT | GTC | GTG | ACA | CAC | TTT | GGG | AAC | CAC | GAA | GAT | GAC | CTC | GAC | AGG |
|------|--------------|-----|-----|-----|-----|-----|-----|-----|-----|----------------------|-----|-----|----------------|-----|-----|--------------|-----|--------------|
| | K | L | Q | Α | I | Α | V | S | K | L | L | K | S | S | S | N | Q | V |
| 1675 | AAA | CTG | CAG | GCT | ATT | GCT | GTT | TCA | AAA | CTA | CTG | AAA | AGT | AGC | TCT | AAT | CAA | GTG |
| | I | F | L | G | Y | Ī | T | S | A | P | G | S | R | D | Y | \mathbf{L} | Q | \mathbf{L} |
| 1729 | ATA | TTT | CTG | GGA | TAT | ATC | ACT | TCA | GCA | CCT | GGC | TCC | AGA | GAT | TAT | CTA | CAG | CTC |
| | T | E | H | G | N | V | K | D | I | D | S | T | D | Н | D | R | W | C |
| 1783 | ACT | GAA | CAT | GGC | AAT | GTG | AAG | GAT | ATC | GAC | AGC | ACT | \mathtt{GAT} | CAT | GAC | AGA | TGG | TGT |
| | E | Y | I | M | Y | R | G | Ļ | I | R | Ŀ | G | Y | Α | R | I | S | H |
| 1837 | GAA | TAC | ATT | ATG | TAT | CGA | GGG | CTG | ATC | AGG | TTG | GGT | TAT | GCA | AGA | ATC | TCC | CAT |
| | Α | E | L | S | D | S | E | I | Q | M | Α | K | F | R | I | P | D | D |
| 1891 | GCT | GAA | CTG | AGT | GAT | TCA | GAA | ATT | CAG | ATG | GCA | AAA | TTT | AGG | ATC | CCT | GAT | GAC |
| | P | T | N | Y | R | Ď | N | Q | K | V | V | I | D | H | R | E | V | S |
| 1945 | CCC | ACT | AAT | TAT | AGA | GAC | AAC | CAG | AAA | GTG | GTC | ATA | GAC | CAC | AGA | GAA | GTT | TCT |
| | \mathbf{E} | K | I | H | F | N | P | R | F | G | S | Y | K | E | G | Н | N | Y |
| 1999 | GAG | AAA | ATT | CAT | TTT | AAT | CCC | AGA | TTT | GGA | TCC | TAC | AAA | GAA | GGA | CAC | AAT | TAT |
| | E | N | N | H | H | F | H | M | N | T | P | K | Y | F | L | * | | |
| 2053 | GAA | AAC | AAC | CAT | CAT | TTT | CAT | ATG | AAT | ACT | CCC | AAA | TAC | TTT | TTA | TGA | AAC | |

Figure 3 Amino acid sequence of 125P5C8.

| 1 | MTSLWREILL | ESLLGCVSWS | LYHDLGPMIY | YFPLQTLELT | GLEGFSIAFL | 50 |
|-----|-------------|------------|------------|------------|------------|-----|
| 51 | SPIFLTITPF | WKLVNKKWML | TLLRIITIGS | IASFQAPNAK | LRLMVLALGV | 100 |
| 101 | SSSLIVQAVT | WWSGSHLQRY | LRIWGFILGQ | IVLVVLRIWY | TSLNPIWSYQ | 150 |
| 151 | MSNKVILTLS | AIATLDRIGT | DGDCSKPEEK | KTGEVATGMA | SRPNWLLAGA | 200 |
| 201 | AFGSLVFLTH | WVFGEVSLVS | RWAVSGHPHP | GPDPNPFGGA | VLLCLASGLM | 250 |
| 251 | LPSCLWFRGT | GLIWWVTGTA | SAAGLLYLHT | WAAAVSGCVF | AIFTASMWPQ | 300 |
| 301 | TLGHLINSGT | NPGKTMTIAM | IFYLLEIFFC | AWCTAFKFVP | GGVYARERSD | 350 |
| 351 | VLLGTMMLII | GLNMLFGPKK | NLDLLLQTKN | SSKVLFRKSE | KYMKLFLWLL | 400 |
| 401 | VGVGLLGLGL | RHKAYERKLG | KVAPTKEVSA | AIWPFRFGYD | NEGWSSLERS | 450 |
| 451 | AHLLNETGAD | FITILESDAS | KPYMGNNDLT | MWLGEKLGFY | TDFGPSTRYH | 500 |
| 501 | TWGIMALSRY | PIVKSEHHLL | PSPEGEIAPA | ITLTVNISGK | LVDFVVTHFG | 550 |
| 551 | NHEDDLDRKL | QAIAVSKLLK | SSSNQVIFLG | YITSAPGSRD | YLQLTEHGNV | 600 |
| 601 | KDIDSTDHDR | WCEYIMYRGL | IRLGYARISH | AELSDSEIQM | AKFRIPDDPT | 650 |
| 651 | MABDMOKAVIT | DHREVSEKIH | FNDRFGSYKE | CHNYENNHHE | HMNTPKYFL | 699 |

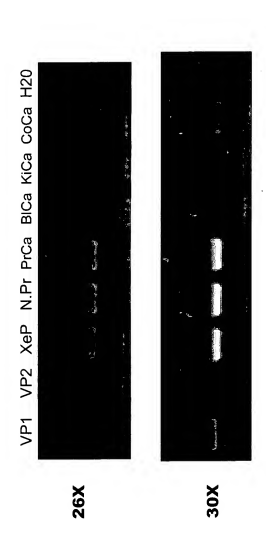
Figure 4A Alignment with AK025164 protein product Score = 1397 bits (3615), Expect = 0.0 Identities = 682/699 (97%), Positives = 683/699 (97%)

| Query: | 1 | MTSLWREILLESLLGCVSWSLYHDLGPMIYYFPLQTLELTGLEGFSIAFLSPIFLTITPF MTSLWREILLESLLGCVSWSLYHDLGPMIYYFPLQTLELTGLEGFSIAFLSPIFLTITPF | 60 |
|--------|-----|---|-----|
| Sbjct: | 1 | MTSLWREILLESLLGCVSWSLYHDLGPMIYYFPLQTLELTGLEGFSIAFLSPIFLTITPF | 60 |
| Query: | 61 | WKLVNKKWMLTLLRIITIGSIASFQAPNAKLRLMVLALGVSSSLIVQAVTWWSGSHLQRY | 120 |
| Sbjct: | 61 | WKLVNKKWMLTLLRIITIGSIASFQAPNAKLRLMVLALGVSSSLIVQAVTWWSGSHLQRY WKLVNKKWMLTLLRIITIGSIASFQAPNAKLRLMVLALGVSSSLIVQAVTWWSGSHLQRY | 120 |
| Query: | 121 | LRIWGFILGQIVLVVLRIWYTSLNPIWSYQMSNKVILTLSAIATLDRIGTDGDCSKPEEK | 180 |
| Shict | 121 | LRIWGFILGQIVLVVLRIWYTSLNPIWSYQMSNKVILTLSAIATLDRIGTDGDCSKPEEK LRIWGFILGQIVLVVLRIWYTSLNPIWSYQMSNKVILTLSAIATLDRIGTDGDCSKPEEK | 180 |
| - | | | |
| Query: | 181 | KTGEVATGMASRPNWLLAGAAFGSLVFLTHWVFGEVSLVSRWAVSGHPHPGPDPNPFGGA KTGEVATGMASRPNWLLAGAAFGSLVFLTHWVFGEVSLVSRWAVSGHPHPGPDPNPFGGA | 240 |
| Sbjct: | 181 | KTGEVATGMASRPNWLLAGAAFGSLVFLTHWVFGEVSLVSRWAVSGHPHPGPDPNPFGGA | 240 |
| Query: | 241 | VLLCLASGLMLPSCLWFRGTGLIWWVTGTASAAGLLYLHTWAAAVSGCVFAIFTASMWPQ | 300 |
| Shict | 241 | VLLCLASGLMLPSCLWFRGTGLIWWVTGTASAAGLLYLHTWAAAVSGCVFAIFTASMWPQ VLLCLASGLMLPSCLWFRGTGLIWWVTGTASAAGLLYLHTWAAAVSGCVFAIFTASMWPQ | 300 |
| • | | | |
| Query: | 301 | TLGHLINSGTNPGKTMTIAMIFYLLEIFFCAWCTAFKFVPGGVYARERSDVLLGTMMLII TLGHLINSGTNPGKTMTIAMIFYLLEIFFCAWCTAFKFVPGGVYARERSDVLLGTMMLII | 360 |
| Sbjct: | 301 | TLGHLINSGTNPGKTMTIAMIFYLLEIFFCAWCTAFKFVPGGVYARERSDVLLGTMMLII TLGHLINSGTNPGKTMTIAMIFYLLEIFFCAWCTAFKFVPGGVYARERSDVLLGTMMLII | 360 |
| Ouerv: | 361 | GLNMLFGPKKNLDLLLQTKNSSKVLFRKSEKYMKXXXXXXXXXXXXXXXXRHKAYERKLG | 420 |
| Quoty. | 551 | GLNMLFGPKKNLDLLLQTKNSSKVLFRKSEKYMK RHKAYERKLG | |
| Sbjct: | 361 | GLNMLFGPKKNLDLLLQTKNSSKVLFRKSEKYMKLFLWLLVGVGLLGLGLRHKAYERKLG | 420 |
| Query: | 421 | KVAPTKEVSAAIWPFRFGYDNEGWSSLERSAHLLNETGADFITILESDASKPYMGNNDLT | 480 |
| Shigt. | 121 | KVAPTKEVSAAIWPFRFGYDNEGWSSLERSAHLLNETGADFITILESDASKPYMGNNDLT KVAPTKEVSAAIWPFRFGYDNEGWSSLERSAHLLNETGADFITILESDASKPYMGNNDLT | 480 |
| _ | | | |
| Query: | 481 | MWLGEKLGFYTDFGPSTRYHTWGIMALSRYPIVKSEHHLLPSPEGEIAPAITLTVNISGK | 540 |
| Sbict: | 481 | MWLGEKLGFYTDFGPSTRYHTWGIMALSRYPIVKSEHHLLPSPEGEIAPAITLTVNISGK MWLGEKLGFYTDFGPSTRYHTWGIMALSRYPIVKSEHHLLPSPEGEIAPAITLTVNISGK | 540 |
| • | | | |
| Query: | 541 | LVDFVVTHFGNHEDDLDRKLQAIAVSKLLKSSSNQVIFLGYITSAPGSRDYLQLTEHGNV- LVDFVVTHFGNHEDDLDRKLQAIAVSKLLKSSSNQVIFLGYITSAPGSRDYLQLTEHGNV | 600 |
| Sbjct: | 541 | LVDFVVTHFGNHEDDLDRKLQAIAVSKLLKSSSNQVIFLGYITSAPGSRDYLQLTEHGNV | 600 |
| Query: | 601 | KDIDSTDHDRWCEYIMYRGLIRLGYARISHAELSDSEIQMAKFRIPDDPTNYRDNQKVVI | 660 |
| | | KDIDSTDHDRWCEYIMYRGLIRLGYARISHAELSDSEIQMAKFRIPDDPTNYRDNQKVVI | |
| Sbjct: | 601 | KDIDSTDHDRWCEYIMYRGLIRLGYARISHAELSDSEIQMAKFRIPDDPTNYRDNQKVVI | 660 |
| Query: | 661 | DHREVSEKIHFNPRFGSYKEGHNYENNHHFHMNTPKYFL 699 | |
| | | DHREVSEKIHFNPRFGSYKEGHNYENNH+FHMNTPKYFL | |
| Sbjct: | 661 | DHREVSEKIHFNPRFGSYKEGHNYENNHNFHMNTPKYFL 699 | |

Figure 4B Alignment with yeast YCR017 cp Protein

Score = 261 bits (668), Expect = 1e-68 Identities = 204/705 (28%), Positives = 330/705 (45%), Gaps = 50/705 (7%) Query: 15 GCVSWSLYHDLGPMIYYFPLQTLELTGLEGFSIAFLSPIFLTITPFWKLVNKKWMLTLLR 74 G + WS L I++FPL + ++G E + +L PIFL + PFSbjct: 279 GFLFWSNVTSLLCSIWHFPLWYMGISGYEAAILGYLGPIFLYL-PFVSEAFTQYGVLLGG 337 Query: 75 IITIGSIASFQAPNAKLRLMVLALGVSSSLIVQAVTWWSGSHLQ-RYLRIWGFILGQIVL 133 Q P +L + + + + + VQ + + + + + W +LG + TT TG+ Sbjct: 338 IIAIGAYI-VQMPELRLISVAVGTSITVATFVQNLRYITNAETSFSFALTW--LLGLVAS 394 Query: 134 VVLRIWYTSLNPIWSYQMS----NKVILTLSAIATLDRIGTDGDCSKPEEKKTGEVATG 188 V+L++ + + NP W NK L L+ + + + + E K+ + + ... Sbjct: 395 VILKMGFYTNNPTWVILDERNGGYNKTALVLTVLFGM--LSPYVNSINFEGKRNAQAKS- 451 Query: 189 MASRPNWLLAGAAFGSLVFLTHWVFGEVSLVSRWAVSGHPHP-GPDPNPFGGAVLLCLAS 247 FGSL+F H + + S WA G+ GP P P+G L C AS L Sbjct: 452 -ASLIGKLFLAVGFGSLLFGIHQLLTDSSTTIYWAWEGYNESHGPLPWPWGA--LTCTVM 508 Query: 248 GLMLPSCLWFRGTGLIWWVTGTASAAGLLY--LHTWAAAV-SGCVFAIFTASMWPQ---T 301 S + F G L + + S A L + W + G + + AI + PSbjct: 509 LFASLSSVKFMGKPLVPCLLLLISTAVLSARSITQWPKYIFGGLLYAIAMLWLVPSYFSA 568 Query: 302 LGHLINSGTNPGKTMTIAMIFYLLEIFFCAWCTAFKFVPGGVYARERSDVLLGTMMLIIG 361 ++ Y++ + W A+ FVP G RE+ + +L I Sbjct: 569 LGQVQNIWV-----YVLSFSVYIIFVLAHVWVVAYAFVPMGWVLREKIETVLAFSSTFII 623 Query: 362 LNMLFGPKKNLDLLLQTKNSSKVLFRKSEKYMKXXXXXXXXXXXXXXXXXXXRHKAYERKLGK 421 N+L+K+F+ T₁ Sbjct: 624 IGALTCKNLNIQLVTMGKKFFIYVF------FFAVALLSLTARFVYDIRPTGI 670 Query: 422 VAP----TKEVSAAIWPFRFGYDNEGWSSLERSAHLLNETGADFITILESDASKPYMGNN 477 ++ ++A IW FG DN+ W+S +R +L+ + D + +LE+D + MGN Sbjct: 671 PQPYHPDSQLITAGIWTIHFGLDNDMWASEDRMINLIKDMELDVVGLLETDTQRITMGNR 730 Query: 478 DLTMWLGEKLGFYTDFGPSTRYHTWGIMALSRYPIVKSEHHLLPSPEGEIAPAITLTV-N 536 DLT L L Y DFGP HTWG + LS++PIV S HHLLPSP GE+APAI T+ Sbjct: 731 DLTSKLAHDLNMYADFGPGPNKHTWGCVLLSKFPIVNSTHHLLPSPVGELAPAIHATLQT 790 Query: 537 ISGKLVDFVVTHFGNHEDDLDRKLQAIAVSKLLKSSSNQVIFLGYITSAPGSRDY-LQLT 595 + LVD V H G ED+ DR+LQ+ ++KL+ +++ I L Y+ PG +Y ++ Sbjct: 791 YNDTLVDVFVFHSGQEEDEEDRRLQSNYMAKLMGNTTRPAILLSYLVVDPGEGNYNTYVS 850 Query: 596 EHGNVKDIDSTDHDRWCEYIMYRGLIRLGYARISHAELSDSEIQMAKFRIPDDPTNYRDN 655 + DID +D DRWCEYI+YRGL R GYAR++ ++D+E+Q+ KF++ + Sbjct: 851 ETSGMHDIDPSDDDRWCEYILYRGLRRTGYARVARGTITDTELQVGKFQVLSEQA-LVEH 909 Query: 656 QKVVIDHREVSEKIHFNPRFGSYKEGHNYENNHHFHM-NTPKYFL 699 + ++ +SE + + +F G E H +H+ + P+Y+L Sbjct: 910 SDSMYEYGHMSEPEYEDMKFPDKFLGEG-ERGHFYHVFDEPRYYL 953

Figure 5



i

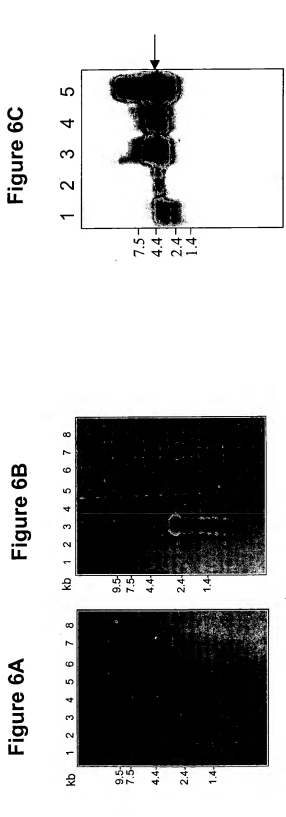


Figure 7

